

O₂-AFFINITY OF FLAVIN RADICAL SPECIES AS STUDIED BY PULSE RADIOLYSIS

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1. Introduction

The activation of molecular oxygen* by metal free biological systems is presently receiving much attention and investigation. For the case of flavins in *aqueous solution*, Massey et al. [1] postulated 'adduct formation' of O₂ and the activating species, reduced flavin (Fl_{red}H₂), (scheme 1, A) in obvious violation of spin conservation. This was based upon kinetic evidence obtained from stopped flow and irrespective of the type of bonding involved between Fl and O₂. Later on, studies of the same group [2,3] with protein bound flavin disclosed spectra of short-lived FlH₂-O₂ intermediates in the catalytic pathway of certain monooxygenases ($\lambda_{\max} \approx 370\text{--}410\text{ nm}$).

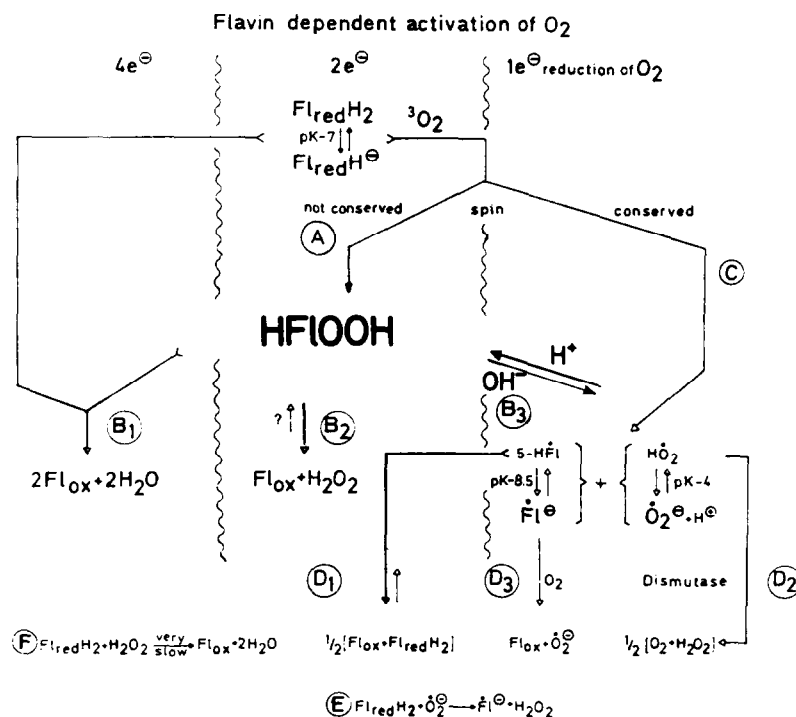
At an earlier stage, Mager and Berends [4,5,6] had provided evidence for an intermediate species in *aprotic* autoxidation of flavin derivatives, which circumvented H₂O₂ formation (scheme 1, B₁). They

proposed for the active intermediate a peroxide structure HFl-OOH. From this it follows that HFl-OOH is an efficient autocatalyst of Fl_{red}H₂ autoxidation under aprotic conditions, while in aqueous solution it decays rapidly as discussed below.

Autocatalysis of Fl_{red}H₂ autoxidation had earlier been found to occur also in aqueous solution [7] and was then thought to be due to the intermediate formation of flavin radical (scheme 1, C). Later on, however, it became obvious from studies of Hemmerich et al. [8], that autocatalysis of Fl_{red}H₂ autoxidation by O₂ is also found under aprotic conditions and in suitably modified flavin systems where – in analogy to the biological case of flavoprotein oxidases – no intermediate radical formation is observed, in agreement with chemical structure.

As next best candidate for the autocatalytic species, superoxide anion $\dot{\text{O}}_2^-$ was demonstrated (step E) by Massey et al. [1] with the aid of superoxide dismutase (step D₂ favored). Stoichiometric amounts of O_2^- , however, could only be detected at pH > 8. The authors suggested this fact as being due to the well known acceleration of superoxide dismutation (pathway D₂). But it is equally obvious that pathway B₃ (scheme 1) is efficiently reversed at pH < 8, i.e. whenever protonation of the radical anion Fl^{•-} ($\text{p}K_{\text{HFl}}^{\text{H}} \approx 8.4$ [9]) becomes

* The use of the term 'O₂-activation' instead of O₂-reduction in the biochemical literature is obviously meant to emphasize the initial polarization of O₂-symmetry and should not be misunderstood in the sense of formation of spectroscopically excited states.



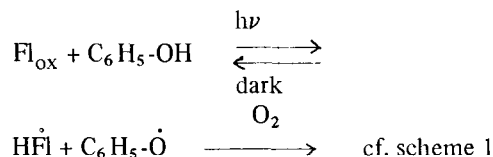
Scheme 1: Pathways of flavin mediated O_2 -reduction. Note that B_1 (unpolar medium) and B_3 (polar medium) are both autocatalytic, the catalysts being HFI-OOH for B_1 and $Fl^{\bullet-}$ (D_3) as well as $O_2^{\bullet-}$ (E) for B_3 . The redox balance is $2 Fl_{ox} + 2 H_2O$ ($= 4e^-$ -transfer) for B_1 , $Fl_{ox} + H_2O_2$ ($= 2e^-$ -transfer) for B_2 and $1/2 Fl_{ox} + O_2^{\bullet-}$ ($= 1e^-$ -transfer) for B_3 , simulating oxygenase (B_1) and e^- -transferase (B_3) action.

predominant. The presence of superoxide dismutase, on the other hand, might force this equilibrium towards the right even at lower pH, thus distorting the whole picture.

Fl_{ox} and H_2O_2 (step F) as final products of the autooxidation can be eliminated from the list of potential catalysts, since extraneous addition of either species did not induce catalysis of $Fl_{red}H_2$ autooxidation in most flavin systems.

Hence, one is tempted to see HFI-OOH as an autocatalytic intermediate by itself and source of 'active oxygen', which could account for three different modes of O_2 -reduction (scheme 1, B_{1-3}), the aprotic medium reflecting flavoprotein oxygenases (B_1), aqueous acid conditions reflecting flavoprotein oxidases (B_2), and alkaline media electron-transferring flavoproteins (B_3). We even suspect a different HFI-OOH isomer being responsible for each of the three reactions [10].

In order to verify this scheme, one of the requirements is the direct measurement of O_2 -affinities inherent in the two flavosemiquinone species that may exist under physiological conditions, i.e. the blue [11,12] HFI and the red [9,12] $Fl^{\bullet-}$. This problem has been tackled first by Vaish and Tollin [13] studying by flash photolysis the flavin-phenol system

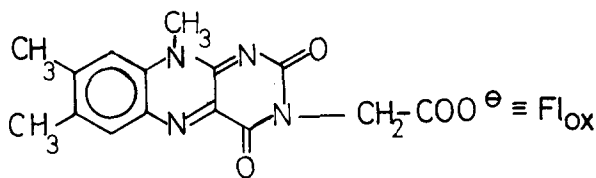


This method has the disadvantage, that the photoreduction of flavin is reversed in the dark and that the presence of phenol and phenoxyl radicals may give rise to disturbing side products and side effects. Hence, we have now used flavin radicals produced by reaction of Fl_{ox}

with hydrated electrons generated by pulse radiolysis [14]. These experiments should decide which of the radical pathways (by way of C or $A \longrightarrow B_3$ (scheme 1)), are essential in flavin dependant O_2 -activation, and over the need to search for mechanisms of rapid intersystem crossing from triplet oxygen other than the 'trivial' but biologically dissatisfactory pathway C.

2. Materials and methods

Lumiflavin-3-acetate



prepared according to Hemmerich [15] was used throughout this study.

The experimental procedure of pulse radiolysis has been described in detail earlier [14,16]. All solutions contained 0.1 M *t*-butanol as scavenger for OH radicals. Saturation by argon, oxygen or air was achieved by the bubbling for ca. 20 min. Various concentrations of O_2 were made up by mixing the appropriate

volumes of argon- and O_2 -saturated solutions. The pH of the solution was maintained either by the lumiflavin acetate or by 10^{-2} M borate or phosphate buffers. Total Fl concentrations varied between $1 \cdot 10^{-5}$ to $5 \cdot 10^{-4}$ M, up to 1% of which were reduced by e_{aq}^- produced by a single pulse of 5 MeV electrons [16].

3. Results

The reaction of hydrated electrons (e_{aq}^-) with Fl_{ox} in anaerobic solution leads to the formation of the radical anion Fl^- in a reaction, which is first order in each reactant, and at a rate of $3 \cdot 10^{10} \text{ M}^{-1} \text{ sec}^{-1}$. The spectrum of the species produced at $4 < \text{pH} < 8$ is that of the neutral radical HFl (fig.1). Since the pK of HFl is 8.4 [9], production of Fl^- is followed by immediate protonation at lower pH. The flavin radicals are found to decay anaerobically in a reaction, which is second order in the decaying species, i.e. by dismutation (fig.2). This reaction is slower for Fl^- because of charge repulsion and reaches at pH 7 a specific rate of $2 \cdot 10^8 \text{ M}^{-1} \text{ sec}^{-1}$. Up to this point our data are essentially an extension of the observations of Land and Swallow [14], who obtained in the anaerobic study $k = 5.7 \times 10^8 \text{ M}^{-1} \text{ sec}^{-1}$ for FMN at pH 5 for the HFl ($\lambda_{\text{max}} = 570 \text{ nm}$) dismutation (D_1 , scheme 1). Our slightly lower values may be due to the fact that our Fl-model (lumiflavin-

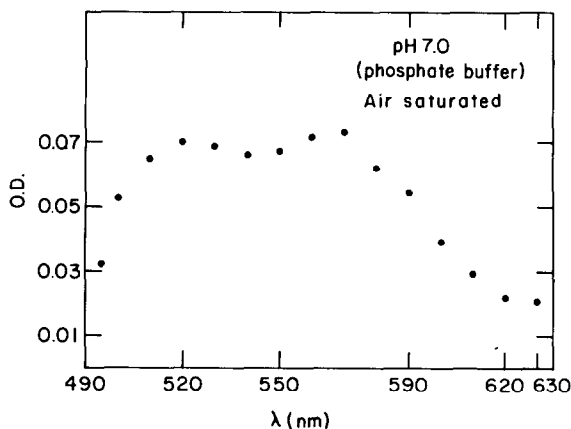
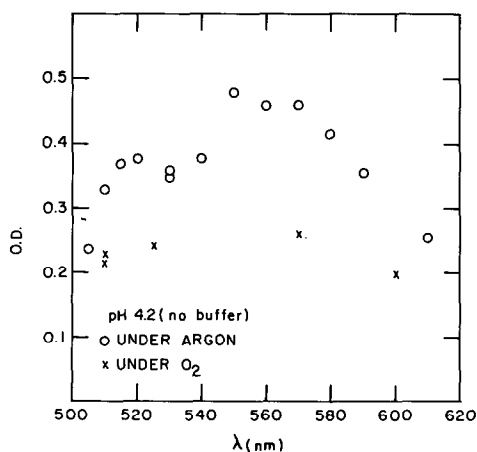


Fig.1. The transient absorption spectra produced by the reaction of e_{aq}^- with lumiflavin-3-acetate under different conditions of pH and saturating gases.

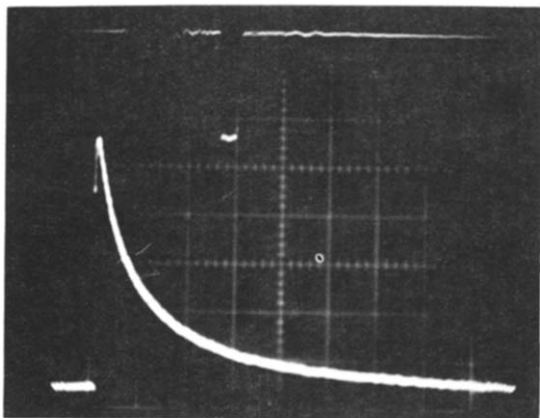


Fig.2. The second order decay of the transient absorption of HFI in aqueous phosphate buffered solution pH 7.0 sweep rate: 500 $\mu\text{sec}/\text{scale unit}$, sensitivity 20 mv/scale unit followed at 570 nm.

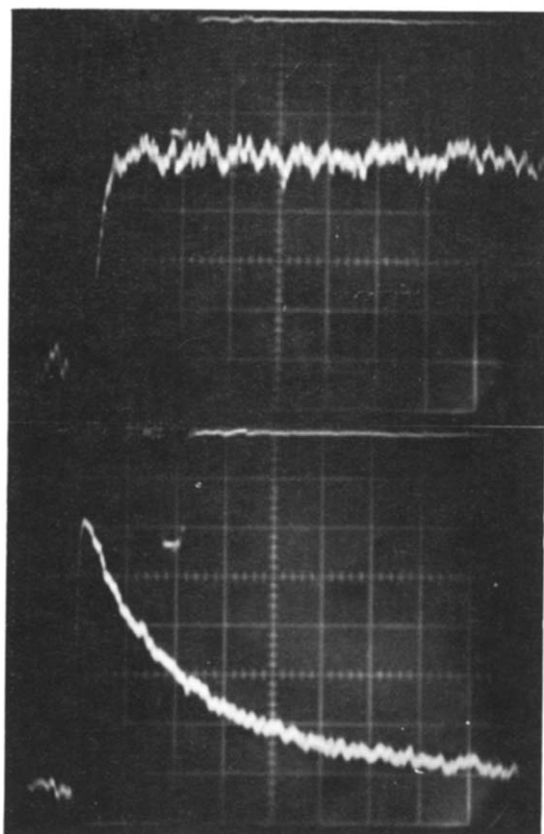


Fig.3. Formation (upper trace) and decay (lower trace) of the HFI radical in air saturated neutral solution followed at 570 nm. Sweep rates: upper 10 $\mu\text{sec}/\text{scale unit}$. Sensitivity 5 mv/scale unit.

3-acetate) carries a negative charge closer to the redox-system, if compared with the phosphate charge of FMN.

In aerobic solution, however, the flavin radical decay becomes strongly pH-dependent: At pH 10, the formation of Fl^- ($\lambda_{\text{max}} = 480 \text{ nm}$) may be observed directly, but only at $[\text{O}_2] \leq 2 \cdot 10^{-4}$. The radical ($[\text{Fl}^-] \approx 5 \cdot 10^{-7}$) is decaying under these conditions in a pseudo first order process at a rate of $2.5 \pm 0.3 \times 10^8 \text{ sec}^{-1} \text{ M}^{-1}$, by reaction with O_2 (D_3 , scheme 1). This is in very good agreement with the value of $3 \times 10^8 \text{ M}^{-1} \text{ sec}^{-1}$ extrapolated for the same reaction by Vaish and Tollin [13] at alkaline pH in their flash photolysis study. At pH 7, however, we observe the formation of HFI in the presence of O_2 -concentrations up to 10^{-3} M . This requires, however, total Fl-concentration to be increased towards 10^{-4} M in order to compete with the reaction of e_{aq}^- with O_2 . The presence of O_2 does not alter the HFI-spectrum and the radical decay pattern remains under these conditions pseudo first order ($k = 2.5 \times 10^7 \text{ M}^{-1} \text{ sec}^{-1}$). At O_2 lower than 10^{-4} M , however, the decay becomes first order in both, HFI and O_2 , at pH 7. The above mentioned anaerobic decay mode, i.e. dismutation, is at pH 7 still negligible, as compared to the autoxidation. If we assume, that the autoxidative decay of HFI is occurring entirely via Fl^- , we can calculate from the rate observed at pH 7 the specific rate for Fl^- -autoxidation, using the expression $k_{\text{spec}} = k_{\text{obs}} / (1 + \text{H}^+/\text{K})$ with $\text{pK} = 8.4$ [9], and we obtain $k_{\text{spec}} = 4.1 \times 10^8 \text{ M}^{-1} \text{ sec}^{-1}$, in good agreement with the above mentioned value, we determined directly at pH 10. Hence, the neglect of HFI as O_2 -reductant is justified. Consequently, the aerobic radical decay changes drastically in shape from pseudo first (i.e. autoxidation) to second order in HFI (i.e. dismutation) upon lowering the pH further. At pH 4.1 the aerobic and anaerobic decay of HFI become identical, $k = 2 \cdot 10^8 \text{ M}^{-1} \text{ sec}^{-1}$.

4. Discussion

It turns out that the two flavin radical species which may exist in the physiological range of pH, HFI and Fl^- , differ greatly in oxygen affinity, in agreement with the results of Vaish and Tollin [13]. With the one exception of glucose oxidase, flavoproteins are known

to stabilize only one of the above mentioned radical species alternatively [10,12]: O_2 -activating flavoproteins produce no superoxide while e^- -transferring flavoproteins — when forced to react with the 'false' acceptor O_2 — produce stoichiometric amounts of it.

We can combine our results with previous knowledge in the following way: In O_2 -activating flavoproteins the reduced state $Fl_{red}H_2$ has, by definition, a high O_2 -affinity. These flavoproteins stabilize red radicals Fl^\cdot , as mentioned above, and Fl^\cdot has a high O_2 -affinity, as we find. Hence, up to 2 moles of O_2^\cdot were to be formed, if radicals were essential in the biocatalysis, but none is found.

In e^- -transferring flavoproteins, on the other hand, O_2 -affinity of $Fl_{red}H_2$ is low. These flavoproteins stabilize blue radicals HFl^\cdot , whose O_2 -affinity is even lower, as we find. Hence, one and only one mole of O_2^\cdot must be found, if radicals were essential, and at the most one mole is indeed found. The low O_2 -reactivity of protein bound HFl groups is not due to its O_2 -inaccessibility of the protein active site, as one might suspect. Hence, the oxidative decay of HFl occurs, preferably, via disproportionation, at least in the free state. But this seems even to be valid for the protein bound HFl , where the intermolecular self contact is generally more difficult, but still not to be excluded. In many cases the shape of the apoprotein might even be such as to favour interflavin contact, be it inter- or, in the case of the many flavoproteins having two flavins per molecule, intramolecular contact [17].

Applied to the pathways of radical formation in scheme 1, the conclusions read as follows:

1) The pH-independent step C would be followed at $pH > 8$ by D_3 , owing to the high O_2 -affinity of Fl^\cdot , rather than by recombination with O_2^\cdot (reversal of B_3), and would yield two moles of O_2^\cdot per Fl_{ox} formed. The actual yield, however, is always less than 1 mole. This rules out pathway C in favour of $A \longrightarrow B_3$, where B_3 still competes with B_1 and B_2 .

2) At $pH < 8$, B_3 is also suppressed, owing to the low O_2 -affinity of HFl^\cdot , thus leaving only B_1 (unpolar) and B_2 (polar) open for O_2 -reduction.

Hence, mechanisms of intersystem crossing of 3O_2 other than O_2^\cdot radical formation should be considered. A proposal has been made elsewhere about how $Fl_{red}H_2$ by itself might facilitate this process [18].

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